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(54) Title: TUMOR ANTIGEN DERIVED GENE-16 (TADG-16): A NOVEL EXTRACÉLLULAR SERINE PROTEASE AND USES THEREOF

(57) Abstract: The present invention provides a DNA encoding a TADG-16 protein selected from the group consisting of: (a) isolated DNA which encodes a TADG-16 protein; (b) isolated DNA which hybridizes to isolated DNA of (a) above and which encodes a TADG-16 protein; and (c) isolated DNA differing from the isolated DNAs of (a) and (b) above in codon sequence due to the degeneracy of the genetic code, and which encodes a TADG-16 protein. Also provided is a vector capable of expressing the DNA of the present invention adapted for expression in a recombinant cell and regulatory elements necessary for expression of the DNA in the cell.

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# TUMOR ANTIGEN DERIVED GENE-16 (TADG-16): A NOVEL EXTRACELLULAR SERINE PROTEASE AND USES THEREOF

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# BACKGROUND OF THE INVENTION

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# Field of the Invention

The present invention relates generally to the fields of cellular biology and the diagnosis of neoplastic disease. More specifically, the present invention relates to an extracellular serine protease termed Tumor Antigen Derived Gene-16 (TADG-16), which is expressed in normal ovaries and testes, as well as certain ovarian carcinomas.

# Description of Related Art

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To date, ovarian cancer remains the number one killer of women with gynecologic malignant hyperplasia. Approximately 75% of women diagnosed with such cancers are already at the high-stage (III and IV) of the disease at their initial diagnosis. During the past 20 years, neither diagnosis nor five year survival have greatly improved for these patients. This is substantially due to the significant number of high-stage initial detections of the disease. Therefore, the challenge remains to develop new markers to improve early diagnosis, and thereby reduce the percentage of high-stage initial diagnoses.

A good tumor marker useful as an indicator of early disease is needed. Extra-cellular proteases have already been implicated in the growth, spread and metastatic progression of many cancers, thereby implying that some extracellular proteases may be candidates for marker of neoplastic development. This is in part due to the ability of malignant cells not only to grow in situ, but to dissociate from the primary tumor and to invade new surfaces (metastasize). The ability to disengage from one tissue and re-engage the surface of another tissue is what results in the morbidity and mortality associated with this disease.

In order for malignant cells to grow, spread or metastasize, they must have the capacity to invade local host tissue, dissociate or shed from the primary tumor, and for metastasis to

occur, enter survive in the bloodstream, plant by invasion into surface of the target organ and establish an environment conducive for new colony growth (including the induction angiogenic and growth factors). During this progression, tissue barriers have to be degraded, including basement membranes and connective tissue. These barriers further include collagen, laminin, proteoglycans and extracellular matrix glycoproteins, such as fibronectin.

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Degradation of these natural barriers, both surrounding the primary tumor and at sites of metastatic invasion, is believed to be brought about by the action of extracellular proteases. Proteases have been classified into four families: serine proteases, metallo-proteases, aspartic proteases and cysteine proteases. Many proteases have been shown to be involved in the human disease process and these enzymes are targets for inhibition by new therapeutic agents.

Certain individual proteases have already been shown to be induced and overexpressed in a diverse group of cancers, and as such, are potential candidates for markers useful for early diagnosis and possibly therapeutic intervention. Examples of proteases, encompassing members of the metallo-proteases, serine proteases, and cysteine proteases, are listed below.

TABLE 1

Protease Expression in Various Cancers

		Gastric	Brain	Breast	<u>Ovarian</u>
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	Serine	uPA	uPA	NES-1NE	S-1
	Proteases	PAI-1 PAI	-1 uPA	uPA	
			tPA		PAI-2
	Cysteine	Cathepsin	<b>B</b> Cathepsin	L Cathepsin	B Cathepsin B
10	Proteases	Cathepsin	L	Cathepsin	L Cathepsin L
15	Metallo- proteases	Collagenas Stromelysin-l	e* Stromelys * Gelatinase	B MMP-9 Gelatinas	e_A
	uPA, Urokinase-ty	pe plasmin	ogen act	iivatoi, tr	PA, Tissue-type
	plasminogen activato	or; PAI-I, Pla	sminogen a	activator 0 i	nhibitors; PAI-2,
	Plasminogen activa	tor inhibito	rs; NES-1	, Normal	epithelial cell-
	specific-1; MMP,	Matrix P me	etallo-prote	așe. *O	verexpressed in
20	gastrointestinal ulc	ers.			

Significantly, there is a good body of evidence supporting the down regulation or inhibition of individual proteases and a subsequent reduction in invasive capacity or malignancy. In work by Clark et al., (Peptides, 14, 1021-8 (1993)) inhibition of in vitro growth of human small cell lung cancer was demonstrated using a general serine protease inhibitor. More recently, Torres-Rosedo et al., (Proc. Natl. Acad. Sci. USA, 90, 7181-7185 (1993)) demonstrated an

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inhibition of patoma tumor cell growth g specific antisense inhibitors for the serine protease hepsin gene. Metastatic potential has also been shown to be reduced using a synthetic inhibitor (batimastat) of metallo-protease in a mouse model with melanoma cells. Powell et al. (Cancer Research, 53, 417-422 (1993)) presented evidence to confirm that the expression of extracellular proteases in relatively non-invasive tumor cells enhances their malignant progression using a tumor-genic, but non-metastatic, prostate cell line. Specifically, Powell et al. demonstrated enhanced metastasis after introducing and expressing the PUMP-1 metallo-protease There is also a body of data to support the notion that expression of cell surface proteases on relatively non-metastatic cell types increases the invasive potential of such cells.

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Extracellular proteases have been directly associated with tumor growth, shedding of tumor cells and invasion of target organs by tumors. Individual classes of proteases are involved in, but not limited to, (a) digestion of stroma surrounding the initial tumor area; (b) digestion of the cellular adhesion molecules to allow dissociation of tumor cells; and (c) invasion of the basement membrane for metastatic growth and the activation of both tumor growth factors and angiogenic factors.

Interfering in the intracellular signal transduction pathways provides mechanisms for numerous therapeutic

applications. While several proteins have een identified that interfere with various signal transduction mechanisms, novel proteins involved in signal transduction pathways are important to provide alternatives for therapy and drug development.

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The prior art is deficient in that the prior art lacks the nucleotide and amino acid sequences corresponding to tumor antigenderived gene 16 (TADG-16). The prior art further lacks effective means of screening to identify proteases, specifically TADG-16, in normal ovaries and testes and certain ovarian expressed carcinomas. The present invention fulfills this longstanding need and desire in the art.

## SUMMARY OF THE INVENTION

This invention describes a new serine protease enzyme.

The TADG-16 enzyme contains the characteristic features of a serine protease, including the conserved catalytic triad (His-Asp-Ser) and a secretion signal sequence. The TADG-16 transcript is present in carcinomas and normal ovarian tissues as well as in normal testes.

Because TADG-16 is secreted and has a potential for extracellular activation, TADG-16 may have a role in normal or aberrant physiological activity of ovary or testes.

provided a DNA encoding a tumor antigen-derived gene (TADG-16) protein, selected from the following: (a) an isolated DNA which encodes a TADG-16 protein; (b) an isolated DNA which hybridizes under high stringency conditions to the isolated DNA of (a) above and which encodes a TADG-16 protein; and (c) an isolated DNA differing from the isolated DNAs of (a) and (b) above in codon sequence due to the degeneracy of the genetic code, and which encodes a TADG-16 protein. The embodiment further includes a vector comprising the TADG-16 DNA and regulatory elements necessary for expression of the DNA in a cell. Additionally embodied is a vector in which the TADG-16 DNA is positioned in reverse orientation relative to the regulatory elements such that TADG-16 antisense mRNA is produced.

In another embodiment of the present invention, there is provided an isolated and purified TADG-16 protein coded for by DNA selected from the following: (a) an isolated DNA which encodes a TADG-16 protein; (b) an isolated DNA which hybridizes under high stringency conditions to isolated DNA of (a) above and which encodes a TADG-16 protein; and (c) an isolated DNA differing from the isolated DNAs of (a) and (b) above in codon sequence due to the degeneracy of the genetic code, and which encodes a TADG-16 protein.

In another embodiment of the sent invention, there is provided a method for detecting TADG-16 mRNA in a sample, comprising the steps of (a) contacting a sample with a probe which is specific for TADG-16; and (b) detecting binding of the probe to TADG-16 mRNA in the sample. In still yet another embodiment of the present invention, there is provided a kit for detecting TADG-16 mRNA, comprising an oligonucleotide probe specific for TADG-16. A label for detection is further embodied in the kit.

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The present invention additionally embodies a method of detecting TADG-16 protein in a sample, comprising the steps of (a) contacting a sample with an antibody which is specific for TADG-16 or a fragment thereof; and (b) detecting binding of the antibody to TADG-16 protein in the sample. Similarly, the present invention also embodies a kit for detecting TADG-16 protein, comprising an antibody specific for TADG-16 protein or a fragment thereof. Means for detection of the antibody is further embodied in the kit.

In another embodiment, the present invention provides an antibody specific for the TADG-16 protein or a fragment thereof.

In yet another embodiment, the present invention provides

20 a method of screening for compounds that inhibit TADG-16,
comprising the steps of (a) contacting a sample comprising TADG-16
protein with a compound; and (b) assaying for TADG-16 protease
activity. Typically, a decrease in the TADG-16 protease activity in the

presence of the compound relative to TADG protease activity in the absence of the compound is indicative of a compound that inhibits TADG-16.

In still yet another embodiment of the present invention, there is provided a method of inhibiting expression of TADG-16 in a cell, comprising the step of (a) introducing a vector into a cell, whereupon expression of the vector produces TADG-16 antisense mRNA in the cell which hybridizes to endogenous TADG-16 mRNA, thereby inhibiting expression of TADG-16 in the cell.

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Further embodied by the present invention, there is provided a method of inhibiting a TADG-16 protein in a cell, comprising the step of (a) introducing an antibody specific for a TADG-16 protein or a fragment thereof into a cell, whereupon binding of the antibody to the TADG-16 protein inhibits the TADG-16 protein.

In another embodiment of the present invention, there is provided a method of targeted therapy to an individual, comprising the step of (a) administering a compound containing a targeting moiety and a therapeutic moiety to an individual, wherein the targeting moiety is specific for TADG-16.

In another embodiment of the present invention, there is provided a method of diagnosing cancer in an individual, comprising the steps of (a) obtaining a biological sample from an individual; and (b) detecting TADG-16 in the sample. Typically, the presence of

TADG-16 in the sample is indicative of the processor of carcinoma in the individual and the absence of TADG-16 in the sample is indicative of the absence of carcinoma in the individual.

In another embodiment of the present invention, there is provided a method of vaccinating an individual against TADG-16, comprising the steps of (a) inoculating an individual with a TADG-16 protein or fragment thereof that lacks TADG-16 protease activity. It is intended that inoculation with the TADG-16 protein or fragment thereof elicits an immune response in the individual, thereby vaccinating the individual against TADG-16.

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In another embodiment of the present invention, there is provided an immunogenic composition, comprising an immunogenic fragment of TADG-16 and an appropriate adjuvant.

Other and further aspects, features, and advantages of the present invention will be apparent from the following description of the presently preferred embodiments of the invention given for the purpose of disclosure.

### BRIEF DESCRIPTION OF THE DRAWINGS

So that the matter in which the above-recited features, advantages and objects of the invention, as well as others which will

particular descriptions of the invention briefly summarized above may be had by reference to certain embodiments thereof which are illustrated in the appended drawings. These drawings form a part of the specification. It is to be noted, however, that the appended drawings illustrate preferred embodiments of the invention and therefore are not to be considered limiting in their scope.

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Figure 1 shows an alignment of a portion of the TADG-16 protein sequence (SEQ ID No. 7) with other known proteases (Prom, Protease M (SEQ ID No. 3); Try1, Trypsinogen 1 (SEQ ID No. 4); SCCE, Stratum corneum chymotryptic like enzyme (SEQ ID No. 5); and Heps, Hepsin (SEQ ID No. 6)).

Figure 2 shows Northern blot analysis of multiple human tissues using the radioactively labeled catalytic domain as a probe. The 1.4 Kb TADG-16 transcript is present in normal human testes and in certain ovarian tumors, but is not detectable at significant levels in other tissues examined. Hybridization of mRNA to  $\beta$ -tubulin is shown as an internal control.

Figure 3A shows the nucleotide and predicted amino acid sequence of the original subclone from the WISH cDNA containing the TADG-16 catalytic domain. Figure 3B shows a sequence identified from the EST database (Accession #AA620757) with homology to the TADG-16 catalytic domain (encoding bases 614 to 1129) and

including the intranslated region and poly tail of the TADG-16 transcript.

Figure 4 shows the nucleotide sequence of the TADG-16 acid sequence. The cDNA the predicted amino cDNA and corresponding to TADG-16 contains a Kozak's consensus sequence (boxed nucleotides) for the initiation of translation from which a putative protein of 314 amino acids is encoded. The protein contains a secretion signal sequence (italicized) and the conserved amino acids of the catalytic triad of the serine protease family (circled) in the appropriate context (underlined residues). The cDNA also contains a polyadenylation sequence in the 3' untranslated region (underlined nucleotides).

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Figure 5 shows TADG-16 (and  $\beta$ -tubulin) expression in normal and carcinoma cell lines.

Figure 6 shows TADG-16 expression in normal (N), benign (B), low malignant potential (LMP) tumors and carcinomas (C).

Figure 6A shows quantitative PCR of TADG-16 (250 bp) and internal control, β-tubulin (470 bp). Lanes 1-3, normal ovary (cases 5-7, respectively); Lanes 4-5, benign mucinous adenoma tumor (cases 8 & 11, respectively); Lane 6, serous LMP tumor (case 14); Lanes 7-8, clear cell carcinoma (cases 20 & 21, respectively); Lanes 9-11, serous adenocarcinoma (cases 22, 29 and 32, respectively); Lane 12, endometrioid adenocarcinoma (case 35). Figure 6B shows a graph

of expression TADG-16 in normal ovaries ovarian benign, LMP and carcinoma tumors.

### DETAILED DESCRIPTION OF THE INVENTION

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This invention describes a new serine protease enzyme complementary to the series of proteases already identified and characterized in ovarian carcinoma. The TADG-16 enzyme contains the characteristic features of all serine proteases, including the catalytic triad of His-Asp-Ser and a signal secretion conserved sequence. The transcript for this enzyme is present in carcinomas and normal ovarian tissues as well as in normal testes. Because TADG-16 is secreted and has a potential for extracellular activation, TADG-16 may have a role in normal or aberrant physiological activity (i.e., normal or carcinomatous growth) of ovary or testes. Furthermore, because of the presence of TADG-16 mRNA in normal testes, there is a potential role for TADG-16 in normal testicular function (e.g., sterility).

The TADG-16 cDNA is 1129 base pairs long (SEQ ID No. 1) and encodes a 314 amino acid protein (SEQ ID No. 2).

In accordance with the present invention there may be employed conventional molecular biology, microbiology, and

recombinant A techniques within the soft the art. Such techniques are explained fully in the literature. See, e.g., Maniatis, Fritsch & Sambrook, "Molecular Cloning: A Laboratory Manual (1982); "DNA Cloning: A Practical Approach," Volumes I and II (D.N. Glover ed. 1985); "Oligonucleotide Synthesis" (M.J. Gait ed. 1984); "Nucleic Acid Hybridization" [B.D. Hames & S.J. Higgins eds. (1985)]; "Transcription and Translation" [B.D. Hames & S.J. Higgins eds. (1984)]; "Animal Cell Culture" [R.I. Freshney, ed. (1986)]; "Immobilized Cells And Enzymes" [IRL Press, (1986)]; B. Perbal, "A Practical Guide To Molecular Cloning" (1984).

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Therefore, if appearing herein, the following terms shall have the definitions set out below.

As used herein, the term "cDNA" shall refer to the DNA copy of the mRNA transcript of a gene.

As used herein, the term "derived amino acid sequence" shall mean the amino acid sequence determined by reading the triplet sequence of nucleotide bases in the cDNA.

As used herein the term "screening a library" shall refer to the process of using a labeled probe to check whether, under the appropriate conditions, there is a sequence complementary to the probe present in a particular DNA library. In addition, "screening a library" could be performed by PCR.

As seed herein, the term "PCR" refers to the Polymerase Chain Reaction that is the subject of U.S. Patent Nos. 4,683,195 and 4,683,202 to Mullis, as well as other improvements to the process/technique of PCR now known in the art.

The amino acid described herein are preferred to be in the "L" isomeric form. However, residues in the "D" isomeric form can be substituted for any L-amino acid residue, as long as the desired functional property of immunoglobulin-binding is retained by the polypeptide. NH<sub>2</sub> refers to the free amino group present at the amino terminus of a polypeptide. COOH refers to the free carboxy group present at the carboxy terminus of a polypeptide. In keeping with standard polypeptide nomenclature, *J Biol. Chem.*, 243:3552-59 (1969), abbreviations for amino acid residues are shown in Table 2.

15 **TABLE 2** 

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		_Symbol	Amino acid
	1_Letter	3 Letter	
	· A	Ala	Alanine
	Ċ	Cys	Cysteine
20	D	Asp	Aspartic acid
	E	Glu	Glutamic acid
	F	Phe	Phenylalanine
	G	Gly	Glycine
	H	His	Histidine
25	I	Ile	Isoleucine
•	K	Lys	Lysine
	L	Leu	Leucine
	M	Met	Methionine
	N	Asn	Asparagine

P Q R S S T V W Y	Pro Gln Arg Ser Thr Val Trp Tyr	Proline Glutamine Arginine Serine Threonine Valine Tryptophan Tyrosine
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It should be noted that all amino-acid residue sequences are represented herein by formulae whose left and right orientation is in the conventional direction of amino-terminus to carboxy-terminus. Furthermore, it should be noted that a dash at the beginning or end of an amino acid residue sequence indicates a peptide bond to a further sequence of one or more amino-acid residues. The above table is presented to correlate the three-letter and one-letter notations which may appear alternately herein.

A "replicon" is any genetic element (e.g., plasmid, chromosome, virus) that functions as an autonomous unit of DNA replication in vivo; i.e., capable of replication under its own control.

A "vector" is a replicon, such as plasmid, phage or cosmid, to which another DNA segment may be attached so as to bring about the replication of the attached segment.

A "DNA molecule" refers to the polymeric form of deoxyribonucleotides (adenine, guanine, thymine, or cytosine) in its either single stranded form, or a double-stranded helix. This term refers only to the primary and secondary structure of the molecule,

and does not light it to any particular tertiary rms. Thus, this term includes double-stranded DNA found, inter alia, in linear DNA molecules (e.g., restriction fragments), viruses, plasmids, and chromosomes. The structure is discussed herein according to the normal convention of giving only the 5' to 3' sequence of the nontranscribed strand of DNA (i.e., the strand having a sequence homologous to the mRNA).

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An "origin of replication" refers to those DNA sequences that participate in DNA synthesis.

A DNA "coding sequence" is a double-stranded 10 sequence which is transcribed and translated into a polypeptide in vivo when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon 15 at the 3' (carboxyl) terminus. A coding sequence can include, but is not limited to, prokaryotic sequences, cDNA from eukaryotic mRNA, genomic DNA sequences from eukaryotic (e.g., mammalian) DNA, and synthetic DNA sequences. A polyadenylation signal and transcription termination sequence will usually be located 3' to the 20 coding sequence.

Transcriptional and translational control sequences are

DNA regulatory sequences, such as promoters, enhancers,

polyadenylation gnals, terminators, and the light that provide for the expression of a coding sequence in a host cell.

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A "promoter sequence" is a DNA regulatory region capable of binding RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' to include the minimum number of bases or elements at detectable initiate transcription levels necessary to will be promoter sequence background. Within the initiation site, as well as protein binding domains transcription responsible binding RNA for the of sequences) (consensus Eukaryotic promoters often, but not always, contain polymerase. "TATA" boxes and "CAT" boxes. Prokaryotic promoters contain. Shineconsensus sequences in addition to the -10 and -35 Dalgarno sequences.

An "expression control sequence" is a DNA sequence that controls and regulates the transcription and translation of another DNA sequence. A coding sequence is "under the control" of transcriptional and translational control sequences in a cell when RNA polymerase transcribes the coding sequence into mRNA, which is then translated into the protein encoded by the coding sequence.

A "signal sequence" can be included near the coding sequence. This sequence encodes a signal peptide, N-terminal to the polypeptide, that communicates to the host cell to direct the polypeptide to the cell surface or secrete the polypeptide into the media, and this signal peptide is clipped off by the host cell before the protein leaves the cell. Signal sequences can be found associated with a variety of proteins native to prokaryotes and eukaryotes.

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The term "oligonucleotide", as used herein in referring to the probe of the present invention, is defined as a molecule comprised of two or more ribonucleotides, preferably more than three. Its exact size will depend upon many factors which, in turn, depend upon the ultimate function and use of the oligonucleotide.

The term "primer" used herein as refers a n oligonucleotide, whether occurring naturally as in a restriction digest or produced synthetically, which is capable of acting as a point of initiation of synthesis when placed under conditions in which synthesis of a primer extension product, which complementary to a nucleic acid strand, is induced, i.e., in the presence of nucleotides and an inducing agent such as a DNA polymerase and at a suitable temperature and pH. The primer may be either single-stranded or double-stranded and must be sufficiently long to prime the synthesis of the desired extension product in the presence of the inducing agent. The exact length of the primer will

depend upon my factors, including temperate, source of primer and use the method. For example, for diagnostic applications, depending on the complexity of the target sequence, the oligonucleotide primer typically contains 15-25 or more nucleotides, although it may contain fewer nucleotides.

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The primers herein are selected to be "substantially" to different strands of a particular complementary target DNA that the primers must be sufficiently This means complementary to hybridize with their respective strands. Therefore, the primer sequence need not reflect the exact sequence of the For example, a non-complementary nucleotide fragment template. may be attached to the 5' end of the primer, with the remainder of the primer sequence being complementary to the strand. Alternatively, non-complementary bases or longer sequences can be interspersed into the primer, provided that the primer sequence has sufficient complementary with the sequence or hybridize therewith and thereby form the template for the synthesis of the extension product.

As used herein, the terms "restriction endonucleases" and "restriction enzymes" refer to enzymes, each of which cut double
20 stranded DNA at or near a specific nucleotide sequence.

A cell has been "transformed" by exogenous or heterologous DNA when such DNA has been introduced inside the cell.

The transforming DNA may or may not be integrated (covalently

linked) into the genome of the cell. In P karyotes, yeast, mammalian cells for example, transforming the DNA may be maintained on an episomal element such as a plasmid. With respect to eukaryotic cells, a stably transformed cell is one in which the transforming DNA has become integrated into a chromosome so that it is inherited by daughter cells through chromosome replication. This stability is demonstrated by the ability of the eukaryotic cell to establish cell lines or clones comprised of a population of daughter cells containing the transforming DNA. A "clone" is a population of cells derived from a single cell or ancestor by mitosis. A "cell line" is a clone of a primary cell that is capable of stable growth in vitro for many generations.

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Two DNA sequences are "substantially homologous" when at least about 75% (preferably at least about 80%, and most preferably at least about 90% or 95%) of the nucleotides match over 15 the defined length of the DNA sequences. Sequences that are homologous substantially can be identified by comparing sequences using standard software available in sequence data banks, or in a Southern hybridization experiment under, for example, stringent conditions as defined for that particular system. 20 appropriate hybridization conditions is within the skill of the art. See, e.g., Maniatis et al., supra; DNA Cloning, Vols. I & II, supra; Nucleic Acid Hybridization, supra.

A terologous" region of the NA construct is an identifiable segment of DNA within a larger DNA molecule that is not found in association with the larger molecule in nature. Thus, when the heterologous region encodes a mammalian gene, the gene will usually be flanked by DNA that does not flank the mammalian genomic DNA in the genome of the source organism. In another example, coding sequence is a construct where the coding sequence itself is not found in nature (e.g., a cDNA where the genomic coding sequence contains introns, or synthetic sequences having codons different than the native gene). Allelic variations or naturally-occurring mutational events do not give rise to a heterologous region of DNA as defined herein.

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The labels most commonly employed for these studies are radioactive elements, enzymes, chemicals which fluoresce when exposed to ultraviolet light, and others. A number of fluorescent materials are known and can be utilized as labels. These include, for example, fluorescein, rhodamine, auramine, Texas Red, AMCA blue and Lucifer Yellow. A particular detecting material is anti-rabbit antibody prepared in goats and conjugated with fluorescein through an isothiocyanate.

Proteins can also be labeled with a radioactive element or with an enzyme. The radioactive label can be detected by any of the currently available counting procedures. The preferred isotope may

be selected from <sup>3</sup>H, <sup>14</sup>C, <sup>32</sup>P, <sup>35</sup>S, <sup>36</sup>Cl, <sup>51</sup>Cr, <sup>52</sup>Co, <sup>59</sup>Fe, <sup>90</sup>Y, <sup>125</sup>I, <sup>131</sup>I, and <sup>186</sup>Re.

Enzyme labels are likewise useful, and can be detected by any of the presently utilized colorimetric, spectrophotometric, fluorospectrophotometric, amperometric or gasometric techniques. The enzyme is conjugated to the selected particle by reaction with bridging molecules such as carbodiimides, diisocyanates, glutaraldehyde and the like. Many enzymes which can be used in these procedures are known and can be utilized. The preferred are peroxidase, β-glucuronidase, β-D-glucosidase, β-D-galactosidase, urease, glucose oxidase plus peroxidase and alkaline phosphatase. U.S. Patent Nos. 3,654,090, 3,850,752, and 4,016,043 are referred to by way of example for their disclosure of alternate labeling material and methods.

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A particular assay system developed and utilized in the art is known as a receptor assay. In a receptor assay, the material to be assayed is appropriately labeled and then certain cellular test colonies are inoculated with a quantitiy of both the label after which binding studies are conducted to determine the extent to which the labeled material binds to the cell receptors. In this way, differences in affinity between materials can be ascertained.

An assay useful in the art is known as a "cis/trans" assay.

Briefly, this assay employs two genetic constructs, one of which is

typically a placed that continually expresses articular receptor of interest when transfected into an appropriate cell line, and the second of which is a plasmid that expresses a reporter such as luciferase, under the control of a receptor/ligand complex. Thus, for example, if it is desired to evaluate a compound as a ligand for a particular receptor, one of the plasmids would be a construct that results in expression of the receptor in the chosen cell line, while the second plasmid would possess a promoter linked to the luciferase gene in which the response element to the particular receptor is inserted. If the compound under test is an agonist for the receptor, the ligand will complex with the receptor, and the resulting complex will bind the response element and initiate transcription of the luciferase gene. The resulting chemiluminescence is then measured photometrically, and dose response curves are obtained and compared to those of known ligands. The foregoing protocol is described in detail in U.S. Patent No. 4,981,784.

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As used herein, the term "host" is meant to include not only prokaryotes but also eukaryotes such as yeast, plant and animal cells. A recombinant DNA molecule or gene which encodes a human TADG-16 protein of the present invention can be used to transform a host using any of the techniques commonly known to those of ordinary skill in the art. Especially preferred is the use of a vector containing coding sequences for the gene which encodes a human

TADG-16 protein of the present invention for process of prokaryote transformation. Prokaryotic hosts may include E. coli, S. tymphimurium, Serratia marcescens and Bacillus subtilis. Eukaryotic hosts include yeasts such as Pichia pastoris, mammalian cells and insect cells.

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In general, expression vectors containing promoter sequences which facilitate the efficient transcription of the inserted DNA fragment are used in connection with the host. The expression typically contains an origin of replication, vector promoter(s). terminator(s), as well as specific genes which are capable of providing phenotypic selection in transformed cells. The transformed hosts can be fermented and cultured according to means known in the art to achieve optimal cell growth.

The invention includes a substantially pure DNA encoding a TADG-16 protein, a strand of which DNA will hybridize at high a probe containing a sequence stringency of at least consecutive nucleotides of SEQ ID No. 1. The protein encoded by the DNA of this invention may share at least 80% sequence identity (preferably 85%, more preferably 90%, and most preferably 95%) with the amino acids shown in SEQ ID No. 2. More preferably, the 20 DNA includes the coding sequence of the nucleotides shown in SEQ ID No. 1, or a degenerate variant of such a sequence.

preferably consists of a sequence of at least 20 consecutive nucleotides, more preferably 40 nucleotides, even more preferably 50 nucleotides, and most preferably 100 nucleotides or more (up to 100%) of the coding sequence of the nucleotides shown in SEQ ID No. 1 or the complement thereof. Such a probe is useful for detecting expression of TADG-16 in a cell by a method including the steps of (a) contacting mRNA obtained from the cell with the labeled hybridization probe; and (b) detecting hybridization of the probe with the mRNA.

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This invention also includes a substantially pure DNA containing a sequence of at least 15 consecutive nucleotides (preferably 20, more preferably 30, even more preferably 50, and most preferably all) of the region from nucleotides 1 to 3147 of the nucleotides shown in SEQ ID No. 1.

By "high stringency" is meant DNA hybridization and wash conditions characterized by high temperature and low salt concentration, e.g., wash conditions of 65°C at a salt concentration of approximately 0.1 x SSC, or the functional equivalent thereof. For example, high stringency conditions may include hybridization at about 42°C in the presence of about 50% formamide; a first wash at about 65°C with about 2 x SSC containing 1% SDS; followed by a second wash at about 65°C with about 0.1 x SSC.

substantially pure DNA" is mea DNA that is not part of a milieu in which the DNA naturally occurs, by virtue of separation (partial or total purification) of some or all of the molecules of that milieu, or by virtue of alteration of sequences that flank the claimed DNA. The term therefore includes, for example, a recombinant DNA which is incorporated into a vector, into an autonomously replicating plasmid or virus, or into the genomic DNA of a prokaryote or eukaryote; or which exists as a separate molecule (e.g., a cDNA or a genomic or cDNA fragment produced by polymerase chain reaction (PCR) or restriction endonuclease digestion) independent of other sequences. It also includes a recombinant DNA which is part of a hybrid gene encoding additional polypeptide sequence, e.g., a fusion protein. Also included is a recombinant DNA which includes a portion of the nucleotides shown in SEQ ID No. 1 which encodes an alternative splice variant of TADG-16.

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The DNA may have at least about 70% sequence identity to the coding sequence of the nucleotides shown in SEQ ID No. 1, preferably at least 75% (e.g., at least 80%); and most preferably at least 90%. The identity between two sequences is a direct function of the number of matching or identical positions. When a subunit position in both of the two sequences is occupied by the same monomeric subunit, e.g., if a given position is occupied by an adenine in each of two DNA molecules, then they are identical at that

emple, if 7 positions in a sequence 10 nucleotides in For 4 position. length are identical to the corresponding positions in a second 10nucleotide sequence, then the two sequences have 70% sequence The length of comparison sequences will generally be at identity. least 50 nucleotides, preferably at least 60 nucleotides. 75 nucleotides, and preferably at least most preferably 100 nucleotides. Sequence identity is typically measured using sequence analysis software (e.g., Sequence Analysis Software Package the Genetics Computer Group, University of Wisconsin of Biotechnology Center, 1710 University Avenue, Madison, WI 53705).

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The present invention is directed towards a vector comprising a DNA sequence which encodes a TADG-16 protein, wherein the vector is capable of replication in a host cell, wherein the vector comprises, in operable linkage: a) an origin of replication; b) a promoter; and c) a DNA sequence coding for the TADG-16 protein. Preferably, the vector of the present invention contains a portion of the DNA sequence shown in SEQ ID No. 1.

A "vector" may be defined as a replicable nucleic acid construct, e.g., a plasmid or viral nucleic acid. Vectors may be used to amplify and/or express nucleic acid encoding TADG-16 protein. An expression vector is a replicable construct in which a nucleic acid sequence encoding a polypeptide is operably linked to suitable control sequences capable of effecting expression of the polypeptide

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in a cell. The need for such control sequence will vary depending upon the cell selected and the transformation method Generally, control sequences include a transcriptional enhancer, suitable mRNA ribosomal and/or binding sites, and sequences which control the termination of transcription and translation.

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Methods which are well known to those skilled in the art can be used to construct expression vectors containing appropriate transcriptional and translational control signals. See for example, the techniques described in Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual (2nd Ed.), Cold Spring Harbor Press, N.Y. A gene its transcription control sequences are defined "operably linked" if the transcription control sequences effectively control the transcription of the gene. Vectors of the invention include, but are not limited to, plasmid vectors and viral vectors. Preferred viral vectors of the invention are those derived from retroviruses, adenovirus, adeno-associated virus, SV40 virus, or herpes viruses.

By a "substantially pure protein" is meant a protein which 20 has been separated from at least some of those components which naturally accompany it. Typically, the protein is substantially pure when it is at least 60%, by weight, free from the proteins and other naturally-occurring organic molecules with which it is naturally

Preferably, the purity of the reparation is at least associated in y 75%, more preferably at least 90%, and most preferably at least 99%, by weight. A substantially pure TADG-16 protein may be obtained, for example, by extraction from a natural source; by expression of a recombinant nucleic acid encoding an TADG-16 polypeptide; or by chemically synthesizing the protein. Purity can be measured by any column chromatography such e.g., as appropriate method. using an antibody specific chromatography immunoaffinity TADG-16, polyacrylamide gel electrophoresis, or HPLC analysis. protein is substantially free of naturally associated components when it is separated from at least some of those contaminants which accompany it in its natural state. Thus, a protein which is chemically synthesized or produced in a cellular system different from the cell from which it naturally originates will be, by definition, substantially associated components. Accordingly, from its naturally free substantially pure proteins include eukaryotic proteins synthesized in E. coli, other prokaryotes, or any other organism in which they do not naturally occur.

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In addition to substantially full-length proteins, the invention also includes fragments (e.g., antigenic fragments) of the TADG-16 protein (SEQ ID No. 2). As used herein, "fragment," as applied to a polypeptide, will ordinarily be at least 10 residues, more typically at least 20 residues, and preferably at least 30 (e.g.,

length, but less than the exirc, intact sequence. 50) residues Fragments of the TADG-16 protein can be generated by methods known to those skilled in the art, e.g., by enzymatic digestion of naturally occurring recombinant TADG-16 protein, or bу recombinant DNA techniques using an expression vector that encodes a defined fragment of TADG-16, or by chemical synthesis. The ability of a candidate fragment to exhibit a characteristic of TADG-16 (e.g., binding to an antibody specific for TADG-16) can be assessed by methods described herein. Purified TADG-16 or antigenic fragments of TADG-16 can be used to generate new antibodies or to test existing (e.g., as positive controls in a diagnostic antibodies employing standard protocols known to those skilled in the art.

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Included in this invention are polyclonal antisera generated by using TADG-16 or a fragment of TADG-16 as the immunogen in, e.g., rabbits. Standard protocols for monoclonal and polyclonal antibody production known to those skilled in this art are employed. The monoclonal antibodies generated by this procedure can be screened for the ability to identify recombinant TADG-16 cDNA clones, and to distinguish them from known cDNA clones.

Further included in this invention are TADG-16 proteins which are encoded at least in part by portions of SEQ ID No. 2, e.g., products of alternative mRNA splicing or alternative protein processing events, or in which a section of TADG-16 sequence has

been deleted. fragment, or the intact TAI 16 polypeptide, may be covalently linked to another polypeptide, e.g., which acts as a label, a ligand or a means to increase antigenicity.

The invention also includes a polyclonal or monoclonal antibody which specifically binds to TADG-16. The invention encompasses not only an intact monoclonal antibody, but also an immunologically-active antibody fragment, e.g., a Fab or (Fab)<sub>2</sub> fragment; an engineered single chain Fv molecule; or a chimeric molecule, e.g., an antibody which contains the binding specificity of one antibody, e.g., of murine origin, and the remaining portions of another antibody, e.g., of human origin.

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In one embodiment, the antibody, or a fragment thereof, may be linked to a toxin or to a detectable label, e.g., a radioactive fluorescent isotopic label, non-radioactive label, label, or label, paramagnetic label, enzyme chemiluminescent colorimetric label. Examples of suitable toxins include diphtheria toxin, Pseudomonas exotoxin A, ricin, and cholera toxin. Examples of suitable enzyme labels include malate hydrogenase, staphylococcal nuclease, delta-5-steroid isomerase, alcohol dehydrogenase, alphadehydrogenase, triose phosphate isomerase, glycerol phosphate alkaline phosphatase, asparaginase, glucose oxidase, peroxidase, beta-galactosidase, ribonuclease, urease, catalase, glucose-6phosphate dehydrogenase, glucoamylase, acetylcholinesterase, etc.

Examples of stable radioisotopic labels incl. 3H, 125I, 131I, 32P, 35S, 14C, etc.

Paramagnetic isotopes for purposes of in vivo diagnosis can also be used according to the methods of this invention. There 5 are numerous examples of elements that are useful in magnetic For discussions on in vivo nuclear magnetic resonance imaging. resonance imaging, see, for example, Schaefer et al., (1989) JACC 14, 472-480; Shreve et al., (1986) Magn. Reson. Med. 3, 336-340; Wolf, G. L., (1984) Physiol. Chem. Phys. Med. NMR 16, 93-95; Wesbey et al., 10 (1984) Physiol. Chem. Phys. Med. NMR 16, 145-155; Runge et al., (1984) Invest. Radiol. 19, 408-415. Examples of suitable fluorescent labels include a fluorescein label, an isothiocyalate label. rhodamine label, a phycocrythrin label, a phycocyanin label, an allophycocyanin label, an ophthaldehyde label, a fluorescamine label, etc. Examples of chemiluminescent labels include a luminal label, an 15 isoluminal label, an aromatic acridinium ester label, an imidazole label, an acridinium salt label, an oxalate ester label, a luciferin label, a luciferase label, an aequorin label, etc.

Those of ordinary skill in the art will know of other suitable labels which may be employed in accordance with the present invention. The binding of these labels to antibodies or fragments thereof can be accomplished using standard techniques commonly known to those of ordinary skill in the art. Typical

techniques ar escribed by Kennedy et al., (6) Clin. Chim. Acta 70, 1-31; and Schurs et al., (1977) Clin. Chim. Acta 81, 1-40. Coupling techniques mentioned in the latter are the glutaraldehyde method, the periodate method, the dimaleimide method, the m-maleimidobenzyl-N-hydroxy-succinimide ester method. All of these methods are incorporated by reference herein.

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Also within the invention is a method of detecting TADG-16 protein in a biological sample, which includes the steps of contacting the sample with the labeled antibody, e.g., radioactively tagged antibody specific for TADG-16, and determining whether the antibody binds to a component of the sample.

As described herein, the invention provides a number of diagnostic advantages and uses. For example, the TADG-16 protein is useful in diagnosing cancer in different tissues since this protein is highly overexpressed in tumor cells. Antibodies (or antigen-binding fragments thereof) which bind to an epitope specific for TADG-16, are useful in a method of detecting TADG-16 protein in a biological sample for diagnosis of cancerous or neoplastic transformation. This method includes the steps of obtaining a biological sample (e.g., cells, blood, plasma, tissue, etc.) from a patient suspected of having cancer, contacting the sample with a labeled antibody (e.g., radioactively tagged antibody) specific for TADG-16, and detecting the TADG-16 protein using standard immunoassay techniques such as

an ELISA. Applied binding to the biological ple indicates that the sample contains a component which specifically binds to an epitope within TADG-16.

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Likewise, a standard Northern blot assay can be used to ascertain the relative amounts of TADG-16 mRNA in a cell or tissue obtained from a patient suspected of having cancer, in accordance with conventional Northern hybridization techniques known to those of ordinary skill in the art. This Northern assay uses a hybridization probe, e.g., radiolabelled TADG-16 cDNA, either containing the full-length, single stranded DNA having a sequence complementary to SEQ ID No. 1, or a fragment of that DNA sequence at least 20 (preferably at least 30, more preferably at least 50, and most preferably at least 100) consecutive nucleotides in length. The DNA hybridization probe can be labeled by any of the many different methods known to those skilled in this art.

Antibodies to the TADG-16 protein can be used in an immunoassay to detect increased levels of TADG-16 protein expression in tissues suspected of neoplastic transformation. These same uses can be achieved with Northern blot assays and analyses.

The TADG-16 cDNA is 1129 base pairs long (SEQ ID No. 1) encoding for a 314 amino acid protein (SEQ ID No. 2). The availability of the TADG-16 gene provides numerous utilities. For example, the TADG-16 gene can be used as a diagnostic or therapeutic target in

ovarian and or carcinomas, including bree prostate, lung and colon.

The present invention is directed to DNA encoding a tumor antigen-derived gene (TADG-16) protein, selected from (a) an isolated DNA which encodes a TADG-16 protein; (b) an isolated DNA which hybridizes under high stringency conditions to the isolated DNA of (a) above and which encodes a TADG-16 protein; and (c) an isolated DNA differing from the isolated DNAs of (a) and (b) above in codon sequence due to the degeneracy of the genetic code, and which encodes a TADG-16 protein. It is preferred that the DNA has the sequence shown in SEQ ID No. 1 and the TADG-16 protein has the amino acid sequence shown in SEQ ID No. 2.

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invention is directed toward vector The present comprising the TADG-16 DNA and regulatory elements necessary for expression of the DNA in a cell, or a vector in which the TADG-16 DNA is positioned in reverse orientation relative to the regulatory elements such that TADG-16 antisense mRNA is produced. An antisense molecule corresponding to TADG-16 mRNA is shown in SEQ ID No. 16. The invention is also directed toward host cells transfected with either of the above-described vector(s). Representative host cells are cells, plant cells and insect cells. bacterial cells, mammalian Preferably, the bacterial cell is E. coli.

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The present invention is directed ward an isolated and purified TADG-16 protein coded for by DNA selected from the following: (a) an isolated DNA which encodes a TADG-16 protein; (b) an isolated DNA which hybridizes under high stringency conditions to isolated DNA of (a) above and which encodes a TADG-16 protein; and (c) an isolated DNA differing from the isolated DNAs of (a) and (b) above in codon sequence due to the degeneracy of the genetic code, and which encodes a TADG-16 protein. Preferably, the protein has the amino acid sequence shown in SEQ ID No. 2.

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The present invention is directed toward a method for detecting TADG-16 mRNA in a sample, comprising the steps of (a) contacting a sample with a probe which is specific for TADG-16; and (b) detecting binding of the probe to TADG-16 mRNA in the sample. The present invention is also directed toward a method of detecting TADG-16 protein in a sample, comprising the steps of (a) contacting a sample with an antibody which is specific for TADG-16 or a fragment thereof; and (b) detecting binding of the antibody to TADG-16 protein in the sample. Generally, the sample is a biological sample; preferably, the biological sample is from an individual; and typically, the individual is suspected of having cancer.

The present invention is directed toward a kit for detecting TADG-16 mRNA, comprising an oligonucleotide probe, wherein the probe is specific for TADG-16. The kit may further comprise a label

with which to el the probe; and means for ecting the label. The present invention is additionally directed toward a kit for detecting TADG-16 protein, comprising an antibody which is specific for TADG-16 protein or a fragment thereof. The kit may further comprise means to detect the antibody.

The present invention is directed toward a antibody which is specific for TADG-16 protein or a fragment thereof.

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The present invention is directed toward a method of screening for compounds that inhibit TADG-16, comprising the steps of: (a) contacting a sample containing TADG-16 protein with a compound; and (b) assaying for TADG-16 protease activity. Typically, a decrease in the TADG-16 protease activity in the presence of the compound relative to TADG-16 protease activity in the absence of the compound is indicative of a compound that inhibits TADG-16.

The present invention is directed toward a method of inhibiting expression of TADG-16 in a cell, comprising the step of: (a) introducing a vector expressing TADG-16 antisense mRNA into a cell which hybridizes to endogenous TADG-16 mRNA, thereby inhibiting expression of TADG-16 in the cell. Generally, the inhibition of TADG-16 expression is for treating cancer.

The present invention is directed toward a method of inhibiting a TADG-16 protein in a cell, comprising the step of (a) introducing an antibody specific for a TADG-16 protein or a fragment

thereof into a l which inhibits the TADG-16 otein. Generally, the inhibition of the TADG-16 protein is for treating cancer.

The present invention is directed toward a method of comprising therapy to an individual, step targeted the of (a)administering a compound having a targeting moiety and therapeutic moiety to an individual, wherein the targeting moiety is specific for TADG-16. Representative targeting moiety antibody specific for TADG-16, a ligand that binds TADG-16 or a ligand binding domain of TADG-16, e.g., a CUB domain, an LDLR Likewise, a representative therapeutic domain, etc. radioisotope, a toxin, a chemotherapeutic agent, an immune stimulant or a cytotoxic agent. Typically, the above-described method is useful when the individual suffers from ovarian cancer, breast cancer, lung cancer, prostate cancer, colon cancer or other cancers in which TADG-16 is overexpressed.

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The present invention is directed toward a method of diagnosing cancer in an individual, comprising the steps of (a) obtaining a biological sample from an individual; and (b) detecting TADG-16 in the sample. Generally, the presence of TADG-16 in the sample is indicative of the presence of carcinoma in the individual, and the absence of TADG-16 in the sample is indicative of the absence of carcinoma in the individual. Typically, the biological sample is blood, urine, saliva tears, interstitial fluid, ascites fliud, tumor tissue

TADG-16 are by Northern blot, Western blot, PCR, dot blot, ELIZA sandwich assay, radioimmunoassay, DNA array chips or flow cytometry (after labeling tumor cells). This method may be useful in diagnosing cancers such as ovarian, breast, lung, colon, prostate and others with increased TADG-16 expression.

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The present invention is also directed to an antisense oligonucleotide having the nucleotide sequence complementary to a TADG-16 mRNA sequence. The present invention is also directed to a composition comprising such an antisense oligonucleotide and a physiologically acceptable carrier therefore.

The present invention is also directed to a method of treating a neoplastic state in an individual in need of such treatment, comprising the step of administering to said individual an effective dose of an antisense oligonucleotide. Preferably, the neoplastic state is ovarian cancer, breast cancer and other cancers that exhibit TADG-16 overexpression. For such therapy, the oligonucleotides alone or in combination with other anti-neoplastic agents can be formulated for a variety of modes of administration, including systemic, topical or localized administration. Techniques and formulations generally can be found in *Remington's Pharmaceutical Sciences* (Mack Publishing Co., Easton, PA). The oligonucleotide active ingredient is generally combined with a pharamceutically acceptable carrier such as a

diluent or exciient which can include fill extenders, binders. wetting agents, disintergrants, surface active agents or lubricants, depending on the nature of the mode of administration and dosage Typical dosage forms include tablets, powders, forms. preparations including suspensions, emulsions, and solutions, granules, capsules and suppositories, as well as liquid preparations for injections, including liposome preparations.

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administration, systemic injection For is preferred. including intramuscular, intravenous, intraperitoneal and subcutaneous. For injection, the oligonucleotides of the invention are formulated liquid solutions, preferably in in physiologically buffers. In addition, the oligonucleotides compatible formulated in solid form and redissolved or suspended immediately prior to use. Lyophilized forms are also incldued. Dosages that can be used for systemic administration preferably range from about 0.01 mg/kg to 50 mg/kg administered once or twice per day. However, different dosing schedules can be utilized depending on (1) the potency of an individual oligonucleotide at inhibiting the activity of its target DNA, (2) the severity or extent of the pathological disease state, or (3) the pharmacokinetic behavior of a given oligonucleotide.

The present invention is directed toward a method of vaccinating an individual against TADG-16, comprising the steps of (a) inoculating an individual with a TADG-16 protein or fragment thereof

which lacks TG-16 protease activity. The inoculation with the TADG-16 protein or fragment thereof elicits an immune response in the individual, thereby vaccinating the individual against TADG-16. The vaccination with TADG-16 described herein is intended for an individual who has cancer, is suspected of having cancer or is at risk The present invention is also directed toward an of getting cancer. immunogenic composition, comprising an immunogenic fragment of TADG-16 and an appropriate adjuvant. Generally, the TADG-16 fragment useful for vaccinating an individual consists of a 9-residue fragment up to and including a 20-residue fragment. Preferably, the 9-residue fragments have a sequence such as SEQ ID Nos. 17, 18, 19, 77, 78, 79, 80, 97, 98, 99, 137, 138, 139, 140 or 141. Other TADG-16 useful for vaccinating an individual may be readily determined by an individual having ordinary skill in this art using routine techniques.

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The present invention is further directed to a method of regulating the expression of the TADG-16 protein by designing antisense oligonucleotides directed to the DNA encoding the TADG-16 protein. A person having ordinary skill in this art would be able design such antisense oligonucleotides without undue experimentation.

The following examples are given for the purpose of illustrating various embodiments of the invention and are not meant to limit the present invention in any fashion.

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#### **EXAMPLE 1**

### Cloning of the TADG-16 catalytic domain

Using WISH (an amnion derived cell line) cDNA (ATCC) as

10 a template for PCR with degenerate primers designed to the conserved regions surrounding the invariant histidine and serine residues of the catalytic triad of the serine protease family of proteins, a 498 base pair product was obtained that was similar in particular consensus sequences to other known serine proteases (Figure 1).

The sequences of the degenerate primers used in the initial PCR are as follows:

Serp-S (Sense): 5'-TGGGTIGTIACIGCIGCICA(CT)TG-3' (SEQ ID No. 8); and

Serp-S (Antisense): 5'-A(AG)IGGICCICCI(CG)(TA)(AG)TCICC-3'
20 (SEQ ID No. 9).

Reactions were carried out as described by Underwood et al. (Cancer Res., 59, 4435-9 (1999)).

#### **EXAMPLE 2**

#### Detection of TADG-16 mRNA

Using the radioactively labeled catalytic domain as a probe, Northern blot analysis of multiple human tissues revealed that TADG-16 is highly expressed in normal human testes and in some ovarian tumors, but not detectable at significant levels in other tissues examined (Figure 2). More importantly, Northern analysis showed that the TADG-16 transcript is approximately 1.4 kilobases in length.

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#### EXAMPLE 3

### Sequence analysis of TADG-16

Comparison of the TADG-16 catalytic domain to the EST database identified a homologous sequence (Accession No. AA620757) that overlapped a portion of the TADG-16 catalytic domain clone and also included the 3'-untranslated region and poly (A) tail of the TADG-16 transcript (Figure 3). Comparison of the catalytic domain clone to the GenBank non-redundant database identified a genomic cosmid clone (Accession No. AC005361) homologous to the catalytic domain clone. Using the GRAIL exon identification program available through the National Center for

Biotechnology Information, potential exons engling the 5' portion of the TADG-16 transcript were identified.

EXAMPLE 4

#### Cloning of the TADG-16 cDNA

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PCR primer (T16-F1: 5'-A sense GTCAGGCCGCGGAGAGGAG-3' (SEQ ID No. 10)) was designed to the cDNA predicted by the Grail program and used in conjunction with an antisense primer (T16-R2: 5'-ACTCTGGGCCATCAGCTTCT-3' (SEO ID No. 11)) designed to the overlapping EST that included the polyA+ tail (GenBank Accession No. AA620757 encoding bases 614 to 1129 of Additional antisense primers were utilized in 5'-RACE TADG-16). experiments using a human testes cDNA library as template to identify the 40-most 5' bases. The sequence of the 5'-RACE primers are as follows:

T16-R6: 5'-CGGAGGGATCACTAAGGTCACTATACGT-3' (SEQ ID No. 12); and

T16-R7: 5'-TATACGTTTCAAAGCAGTGCGCCGCCGT-3' (SEQ ID No. 13).

This allowed for the identification of the 1129 bases of the sequence reported herein. Within this 1129 bases, there is a Kozak's consensus

sequence for initiation of translation, open reading frame encoding a 314 amino acid protein and a polyadenylation signal.

EXAMPLE 5

Tissue-specific expression of TADG-16

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Using a previously authenticated semi-quantitative PCR technique (Shigemasa et al., *J. Soc. Gynecological Inv.*, 4, 95-102 (1997)), the expression level of the TADG-16 transcript was examined in normal ovarian tissue and ovarian tumor specimens. To do this, a TADG-16-specific PCR product was co-amplified with a PCR product for β-tubulin as an internal control. To amplify a 237 bp PCR product specific for TADG-16, the following primers were used:

T16-F2: 5'-GGTCGCCATCATAAACAACT-3' (SEQ ID No. 14); and T16-R2: 5'-ACTCTGGGCCATCAGCTTCT-3' (SEQ ID No. 15).

The reaction mixture was heated to 94°C for 1.5 min, then 30 cycles of PCR was carried out under the following conditions: 30 sec of denaturation at 94°C, 30 sec of annealing at 62°C and 30 sec of extension at 72°C. A final extension at 72°C was performed for 7 min before the reaction was terminated. These PCR products were electrophoresed through an agarose gel to separate them based on

size. Based ominis experiment, TADG-16 appress to be expressed in tumor tissue (Figures 5 & 6).

**EXAMPLE 6** 

### Expression of TADG-16 in tumors

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The expression of the serine protease TADG-16 gene in low malignant potential tumors, and carcinoma (both and serous type) by quantitative PCR using TADG-16mucinous specific primers was determined (primers directed toward the βtubulin message were used as an internal standard). These data confirm the overexpression of the TADG-16 surface protease gene in ovarian carcinoma, including both low malignant potential tumors and Expression of TADG-16 is increased over normal 1.5 overt carcinoma. levels in low malignant potential tumors, and high stage tumors (Stage III) of this group have higher expression of TADG-16 when compared to low stage tumors (Stage 1) (Table 3). In overt carcinoma, serous tumors exhibit the highest levels of TADG-16 expression, mucinous tumors express levels of TADG-16 comparable with the high stage low malignant potential group.

	Expression	of TADG	i-16			
	Case No.	Code	Stage	Grade	Histology	TADG-16
	1	1	-	_	-	0.553
5	2	1	-	_	-	0.232
	3	1	-	-	<u>.</u>	0.229
	4	1	-	-	· <b>-</b>	0.400
	5	1	-	-	-	0.226
	6	1	-	-	-	0.230
10	7	1	-	-	<del>-</del>	0.269
	. 8	2	-	~	-	0.121
	9	2	-	-	-	0.514
	10	2	-	-	-	0.333
	1 1	2	-	<b>-</b>	-	0.323
15	1 2	3				0.732
	1 3	3			•	0.487
	1 4	3				0.850
	15	4	1	1	2	0.815
	16	4	1	1	3	0.287
20	17	4	1	2	2	0.382
	18	4	1	1	1	0.400
	19	4	1	1	2	0.548
	20	4	1	2	4	2.120
	2 1	4	1	2	4	1.700
25	22	, <b>4</b>	1	1	1	1.760
	2 3	4	1	2	1	1.240
	2 4	4	2	3	1	1.320
	2 5	4	2	1	1	0.710
	26	4	3	1	2	0.828
30	27	4	3	1	1	1.730
	28	4	3	1	i	0.510
	29	4	. 3	1	1	2.320
	3 0	4	3	1	2 3	0.792
	3 1	4	3	1		0.899
35	32	4	3	2	1	1.880
	3 3	4	3	2	1	1.130
	34	4	3	2	3 3	0.892
	35	4	3	2	3	1.990
	36	4	3	2		0.365
40	37	4	3 3 3 3 3 3 3 3 3 3	2 2 2 2 3 3	1	1.840
	38	4	3		1 3	1.430 0.830
	39	4	3	3		
	40	4	3	1	1	1.730
	41 .	4	<b>5</b> .	. 1	1	2.910

Code: 1, normal ovary; 2, benign tumor (adence); 3, LMP tumor; 4, cancer (adence arcinoma).

Stage = Clinical stage: 1, stage 1; 2, stage 2; 3, stage 3.

Grade = Histological grade: 1, grade 1; 2, grade 2; 3, grade 3.

5 Histology: 1, serous carcinoma; 2, mucinous carcinoma; 3, endometrioid carcinoma; 4, clear cell carcinoma.

10 **TABLE 4** 

mRNA Expression	mRNA Expression Levels of TADG-16 Gene in Ovarian Cancers					
_	N mRNA Expression Levels					
		mean	SD			
15						
Normal ovary	7	0.306	0.126			
Benign tumor	4	0.323	0.161			
LMP tumor	3	0.690	0.185			
Ovarian cancer	27	1.235	0.692			
20			•			
Clinical stage		•				
Stage 1	9	1.028	0.695			
Stage 2	2	1.015	0.431			
Stage 3	16	1.380	0.711			
25			· .			
Histological grad	le					
Grade 1	1 4	1.160	0.794			
Grade 2	9	1.300	0.667			
Grade 3	4	1.355	0.415			
30						
Histological type	:					
Serous	1 4	1.494	0.688			
Mucinous	5 .	0.673	0.199			
Endometrio	id 6	0.877	0.609			
35 Clear Cell	2	1.910	0.297			

### TABLE 5

5		p-value <u>(unpaired <i>t</i>-test)</u>
	Tumor type	
	normal vs. benign	0.8473
	normal vs. LMP	0.0046
10	normal vs. cancer	0.0014
	benign vs. LMP	0.0375
	benign vs. cancer	0.0148
	LMP vs. cancer	0.1905
15	Stage	٠.
	stage 1 vs. stage 2	0.9808
	stage 1 vs. stage 3	0.2435
	stage 2 vs. stage 3	0.4951
20	Grade	
	grade 1 vs. grade 2	0.6659
	grade 1 vs. grade 3	0.6472
	grade 2 vs. grade 3	0.8830
25	Histology	<u>.</u>
	serous vs. mucinous	0.0192
	serous vs. endometrioid	0.0743
	serous vs. clear cell	0.4230
	mucinous vs. endometrioid	0.4937
30	mucinous vs. clear cell	0.0012
	endometrioid vs. clear cell	0.0678

### EXAMPLE 7

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## Antisense TADG-16

TADG-16 is cloned and expressed in the opposite orientation such that an antisense RNA molecule (SEQ ID No. 16) is produced. For example, the antisense RNA is used to hybridize to the

TADG-16 RNA into protein.

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#### **EXAMPLE 8**

#### Peptide ranking

For vaccine or immune stimulation, individual 9-mers to 11-mers of the TADG-16 protein were examined to rank the binding of individual peptides to the top 8 haplotypes in the general population (Parker et al., (1994)). The computer program used for this analyses can be found at <a href="http://www-bimas.dcrt.nih.gov/molbio/hla\_bind/">http://www-bimas.dcrt.nih.gov/molbio/hla\_bind/</a>. Table 6 shows the peptide ranking based upon the predicted half-life of each peptide's binding to a particular HLA allele. A larger half-life indicates a stronger association with that peptide and the particular HLA molecule. The TADG-16 peptides that strongly bind to an HLA allele are putative immunogens, and are used to innoculate an individual against hepsin.

## TABLE 6

# TADG-16 peptide ranking

5	HLA Type & Ranking	Start	Peptide	Predicted Dissociation <sub>1/2</sub>	SEQ ID_No.
	HLA A0201				
	1	70	SLLSHRWAL	592.807	17
10	2	299	LLFFPLLWA	395.296	18
	3	142	KLSAPVTYT	329.937	19
	4	96	WMVQFGQLT	94.077	20
	5	10	ALLLARAGL	79.041	2 1
	6	252	QIGVVSWGV	71.726	22
15	7	248	GLWYQIGVV	70.769	23
•	8	139	ALVKLSAPV	69.552	24
	9	291	SQPDPSWPL	66.602	25
	10	130	YLGNSPYDI	47.991	26
	11	190	TLQEVQVAI	42.774	27
20	12	6	ALLLALLLA	42.278	28
	13	165	FENRTDCWV	34.216	29
	1 4	71	LLSHRWALT	21.536	30
•	15	8	LLALLLARA	19.425	31
	16	297	WPLLFFPLL	1.7.136	3 2
25	17	113	QAYYTRYFV	17.002	3 3
	18	123	NIYLSPRYL	10.339	3 4
	19	104	TSMPSFWSL	7.352	
	3 5				
	20	273	NISHHFEWI	7.345	
30	36				
	HLA A0205		·		
	1	70	SLLSHRWAL	25.200	37
	2	4 2	IVGGEDAEL	23.800	38
	3	10	ALLLARAGL	21.000	39
35	4	291	SQPDPSWPL	20.160	40
	5	297	WPLLFFPLL	12.600	4 1
	. 6	248	GLWYQIGVV	12.000	42
	7	8 2	HCFETYSDL	6.300	43

	8	142	KLSAPVTYT	<b>60</b> 00	44
	9	96	WMVQFGQLT	00	45
	10	299	LLFFPLLWA	5.100	46
	1 1	303	PLLWALPLL	4.200	47
5	1 2	123	NIYLSPRYL	4.200	48
	1 3	98	VQFGQLTSM	4.080	49
	1 4	306	WALPLLGPV	3.600	50
	15	71	LLSHRWALT	3.400	5 1
1.0	16	53	WPWQGSLRL	3.150	52
10	17	302	FPLLWALPL	3.150	53
	18	130	YLGNSPYDI	3.000	54
	19	6 190	ALLLALLLA	3.000	5 5
	20	190	TLQEVQVAI	3.000	56
•	HLA A1	•			
15	1	4 4	GGEDAELGR	11.250	57
	2	90	LSDPSGWMV	7.500	58
	3	143	LSAPVTYTK	6.000	59
	4	292	QPDPSWPLL	2.500	60
	5	203	MCNHLFLKY	2.500	61
20	6	87	YSDLSDPSG	1.500	62
	7	168	RTDCWVTGW	1.250	63
	8	47	DAELGRWPW	0.900	6.4
	9	23	SQEAAPLSG	0.675	65
25	10	7	LLLALLLAR	0.500	66
25	11	157	CLQASTFEF	0.500	67
	12	202	SMCNHLFLK	0.500	68
	13 14	111 125	SLQAYYTRY	0.500 0.500	69 70
	15	152	YLSPRYLGN HIQPICLQA	0.500	71
30	16	79	TAAHCFETY	0.500	72
30	17	238	SGGPLACNK	0.500	73
	18	172	WVTGWGYIK	0.400	74
	19	110	WSLQAYYTR	0.300	7.5
	20	191	LQEVQVAII	0.270	76
35	HLA A24	110			
	1	118	RYFVSNIYL	400.000	77
	2	177	GYIKEDEAL	300.000	78
	3	210	KYSFRKDIF	140.000	79
40	4	270	VYTNISHHF	60.000	80
40	5	148	TYTKHIQPI	28.800	8 1
	6 7	300	LFFPLLWAL	24.000	82
	8	234 135	CFGDSGGPL	22.000	83
	٥	133	PYDIALVKL	9.600	84

	9	4	RGALLLALL	40	8.5
	10	<b>1</b> 04	TSMPSFWSL	8.640	86
	11	296	SWPLLFFPL	7.500	87
	1 2	250	WYQIGVVSW	7.200	88
. 5	13	5	GALLLALLL	7.200	89
. <i>5</i>	14	9 5	GWMVQFGQL	7.200	90
	15	199	INNSMCNHL	7.200	91
	16	297	WPLLFFPLL	7.200	92
	17	291	WQPDPSWPL	7.200	93
10	1 8	183	EALPSPHTL	7.200	94
10	19	86	TYSDLSDPS	7.200	95
	20	10	ALLLARAGL	6.000	96
	20	10	ADDDAMAGD	0.000	
	HLA B7				
	1	297	WPLLFFPLL	80.000	97
15	2 3	302	FPLLWALPL	80.000	98
	3	53	WPWQGSLRL	80.000	99
	4	292	QPDPSWPLL	24.000	100
	5	145	APVTYTKHI	24.000	101
	6	42	IVGGEDAEL	20.000	102
20	7	10	ALLLARAGL	18.000	103
	8	104	TSMPSFWSL	12.000	104
	9	183	EALPSPHTL	12.000	105
	10	201	NSMCNHLFL	12.000	106
	1 1	5	GALLLALLL	12.000	107
25	1 2	291	SQPDPSWPL	6.000	108
	1 3	. 70	SLLSHRWAL	6.000	109
	1 4	195	QVAIINNSM	5.000	110
•	1 5	116	YTRYFVSNI	4.000	111
	16	199	INNSMCNHL	4.000	112
30	1 7	8 2	HCFETYSDL	4.000	113
	18	132	GNSPYDIAL	4.000	114
	19	1	MGARGALLL	4.000	115
	20	63	DSHVCGVSL	4.000	116
	HLA B8				
35	1	183	EALPSPHTL	1.600	117
-	2	58	SLRLWDSHV	1.200	118
	3	8 2	HCFETYSDL	1.200	119
	4	116	YTRYFVSNI	1.000	120
	5	2	GARGALLLA	0.800	121
40	6	302	FPLLWALPL	0.800	122
	7	53	WPWQGSLRL	0.800	123
	8	3 1	GPCGRRVIT	0.800	124
	9	297	WPLLFFPLL	0.800	125

	10	5	GALLLALLL	00	126
	1 1	7 1	LLSHRWALT	400	127
	12	242	LACNKNGLW	0.400	128
	13	10	ALLLARAGL	0.400	129
5	1 4	7 0	SLLSHRWAL	0.400	130
	15	63	DŚHVCGVSL	0.400	131
	16	8 9	DLSDPSGWM	0.300	132
	17	132	GNSPYDIAL	0.200	133
	18	140	LVKLSAPVT	0.200	134
10	19	149	YTKHIQPIC	0.200	135
	20	1 5	RAGLRKPES	0.200	136
	HLA B2702				
	1	117.	GRWPWQVSL	1000.000	137
	2	5 1	LRSDQEPLY	300.00	138
15	3	263	RRKLPVDRI	200.000	139
	4	74	SRWRVFAGA	100.000	. 140
	5	128	GRDTSLGRW	100.000	141
	6	266	WRLCGIVSW	60.000	142
	7	3	LRYDGAHLC	60.000	143
20	8	34 .	LRALTHSEL	60.000	144
	9	213	FREWIFQAI	20.000	145
	10	18	GRLPHTQRL	20.000	146
	1 1	101	ERNRVLSRW	20.000	147
•	12	227	NRVLSRWRV	20.000	148
25	1 3	59	SRPKVAALT	20.000	149
	1 4	4 0	VRTAGANGT	20.000	150
	1 5	3 5	QRLLEVISV	18.000	151
	16	98	CQGDSGGPF	10.000	152
	17	112	ARLMVFDKT	6.000	153
30	18	291	WRVFAGAVA	6.000	154
	19	191	GRFLAAICQ	6.000	155
	20	157	CLQASTFEF	3.000	156
	HLA B4403		•		
	. 1	122	SNIYLSPRY	30.000	157
35	2 3	182	DEALPSPHT	24.000	158
		4 5	GEDAELGRW	18.000	159
	4	136	YDIALVKLS	11.250	160
	5	170	DCWVTGWGY	9.000	161
	6	243	ACNKNGLWY	6.000	162
40	7	163	FEFENRTDC	6.000	163
	.8 9	8 8	SDLSDPSGW	6.000	164
	=	79	TAAHCFETY	6.000	165
	10	278	FEWIQKLMA	6.000	166

	1 1	192	QEVQVAIIN	00	167
	12	92	DPSGWMVQF	4.300	168
	13	294	DPSWPLLFF	4.500	169
	14	203	MCNHLFLKY	4.500	170
5	15	76	WALTAAHCF	4.500	171
	16	165	FENRTDCWV	4.000	172
	17	215	KDIFGDMVC	2.500	173
	18	4 8	AELGRWPWQ	2.400	174
	19	272	TNISHHFEW	2.250	175
10	20	227	AQGGKDACF	2.250	176

### **Implications**

That TADG-16 is found at low levels in some normal tissues may not detract from it's potential usefulness as a tumor marker, as there may be an aberrant expression pattern at the translational level that, e.g., allows for detection of TADG-16 in tumor patients but not in healthy patients, and/or activation of the TADG-16 enzyme may be necessary for tumor progression. In the case of the serine protease hepsin, Torres-Rosada et al. demonstrated by down-regulating hepsin that hepsin was required for growth of certain mammalian cells in culture.

The TADG-16 protein sequence is 314 amino acids in length and contains a secretion signal sequence, which suggests that this protein is functional in an extracellular capacity. A proteolytic cleavage site usually associated with protease enzyme activation is present downstream from the secretion signal sequence between

amino acid indues 19 and 20. Moreove the identified clone contains the necessary amino acids characteristic of a functional serine protease catalytic triad, thereby suggesting that this protein may be functioning in a manner that would promote cellular growth or expansion.

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Any patents or publications mentioned in this specification are indicative of the levels of those skilled in the art to which the invention pertains. These patents and publications are herein incorporated by reference to the same extent as if each individual publication was specifically and individually indicated to be incorporated by reference.

One skilled in the art will readily appreciate that the present invention is well adapted to carry out the objects and obtain the ends and advantages mentioned, as well as those inherent therein. examples The present along with the methods, procedures, treatments, molecules, and specific compounds described herein are presently representative of preferred embodiments, are exemplary, and are not intended as limitations on the scope of the invention. Changes therein and other uses will occur to those skilled in the art which are encompassed within the spirit of the invention as defined by the scope of the claims.

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### WHAT IS CLAMED IS:

- DNA encoding a tumor antigen-derived gene (TADG protein, selected from the group consisting of:
  - (a) isolated DNA which encodes a TADG-16 protein;
- (b) isolated DNA which hybridizes under high stringency conditions to the isolated DNA of (a) above and which encodes a TADG-16 protein; and
- (c) isolated DNA differing from the isolated DNAs of (a)
  and (b) above in codon sequence due to the degeneracy of the genetic
  code, and which encodes a TADG-16 protein.
- 2. The DNA of claim 1, wherein said DNA has the sequence shown in SEQ ID No. 1.
  - 3. The DNA of claim 1, wherein said TADG-16 protein has the amino acid sequence shown in SEQ ID No. 2.

4. An oligonucleotide having the nucleotide sequence complementary to a sequence of claim 1.

5 A composition comprising the oligonucleotide according to claim 4 and a physiologically acceptable carrier therefore.

5

- 6. A vector comprising the DNA of claim 1 and regulatory elements necessary for expression of said DNA in a cell.
- 7. The vector of claim 6, wherein said DNA encodes a TADG-16 protein having the amino acid sequence shown in SEQ ID No. 2.
- 8. The vector of claim 6, wherein said DNA is positioned in reverse orientation relative to said regulatory elements such that TADG-16 antisense mRNA is produced.
  - 9. A host cell transfected with the vector of claim 6 said vector expressing a TADG-16 protein.

The host cell of claim 9, where said cell is selected from the group consisting of bacterial cells, mammalian cells, plant cells and insect cells.

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- 11. The host cell of claim 10, wherein said bacterial cell is E. coli.
- 12. Isolated and purified TADG-16 protein coded for by DNA selected from the group consisting of:
  - (a) isolated DNA which encodes a TADG-16 protein;
  - (b) isolated DNA which hybridizes under high stringency conditions to isolated DNA of (a) above and which encodes a TADG-16 protein; and
    - (c) isolated DNA differing from the isolated DNAs of (a) and (b) above in codon sequence due to the degeneracy of the genetic code, and which encodes a TADG-16 protein.
- 13. The TADG-16 protein of claim 12, wherein said 20 protein has the amino acid sequence shown in SEQ ID No. 2.

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14 An antibody, wherein said tibody is specific for TADG-16 protein or a fragment thereof.

- 15. A method for detecting TADG-16 mRNA in a sample, comprising the steps of:
- (a) contacting a sample with a probe, wherein said probe is specific for TADG-16; and
- (b) detecting binding of said probe to TADG-16 mRNA in 10 said sample.
  - 16. The method of claim 15, wherein said sample is a biological sample.
  - 17. The method of claim 16, wherein said biological

sample is from an individual.

18. The method of claim 17, wherein said individual is suspected of having cancer.

19 A kit for detecting TADG-16 RNA, comprising:
an oligonucleotide probe, wherein said probe is specific for TADG-16.

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20. The kit of claim 19, further comprising: a label with which to label said probe; and means for detecting said label.

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- 21. A method of detecting TADG-16 protein in a sample, comprising the steps of:
- (a) contacting a sample with an antibody, wherein said antibody is specific for TADG-16 or a fragment thereof; and
- (b) detecting binding of said antibody to TADG-16 protein in said sample.
- 22. The method of claim 21, wherein said sample is a 20 biological sample.

The method of claim 22, therein said biological sample is from an individual.

- 5 24. The method of claim 23, wherein said individual is suspected of having cancer.
- 25. A kit for detecting TADG-16 protein, comprising:

  an antibody, wherein said antibody is specific for TADG-16

  protein or a fragment thereof.
  - 26. The kit of claim 25, further comprising: means to detect said antibody.

- 27. A method of inhibiting endogenous expression of TADG-16 in a cell, comprising the step of:
- 20 (a) introducing the vector of claim 8 into a cell, wherein expression of said vector produces TADG-16 antisense mRNA in said cell, wherein said TADG-16 antisense mRNA hybridizes to endogenous

TADG-16 mR thereby inhibiting endogeno expression of TADG-16 in said cell.

28. A method of inhibiting a TADG-16 protein in a cell, comprising the step of:

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introducing an antibody into a cell, wherein said antibody is specific for a TADG-16 protein or a fragment thereof, wherein binding of said antibody to said TADG-16 protein inhibits said TADG-16 protein.

- A method of treating a neoplastic state in an 29. individual in need of such treatment, comprising the step individual effective dose the said an 15 administering to oligonucleotide of claim 4.
- 30. The method of claim 29, wherein said neoplastic state is selected from the group consisting of ovarian cancer, breast cancer, lung cancer, colon cancer and prostate cancer.

A method of vaccinating an ividual against TADG-16, comprising the steps of:

inoculating an individual with a TADG-16 protein or fragment thereof, wherein said TADG-16 protein or fragment thereof lack TADG-16 protease activity, wherein said inoculation with said TADG-16 protein or fragment thereof elicits an immune response in said individual, thereby vaccinating said individual against TADG-16.

- 32. The method of claim 31, wherein said TADG-16 fragment is selected from the group consisting of a 9-residue fragment up to a 20-residue fragment.
- 15 33. The method of claim 32, wherein said 9-residue fragment is selected from the group consisting of SEQ ID Nos. 17, 18, 19, 77, 78, 79, 80, 97, 98, 99, 137, 138, 139, 140 and 141.
- 20 34. The method of claim 31, wherein said individual has cancer, is suspected of having cancer or is at risk of getting cancer.

35 An immunogenic composion, comprising an immunogenic fragment of a TADG-16 protein and an adjuvant.

- 36. The immunogenic composition of claim 35, wherein said fragment is selected from the group consisting of a 9-residue fragment up to a 20-residue fragment.
- 37. The immunogenic composition of claim 36, wherein said 9-residue fragment is selected from the group consisting of SEQ ID Nos. 17, 18, 19, 77, 78, 79, 80, 97, 98, 99, 137, 138, 139, 140 and 141.

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- 38. A method of diagnosing cancer in an individual, comprising the steps of:
  - (a) obtaining a biological sample from an individual;
- (b) detecting TADG-16 in said sample, wherein the presence of TADG-16 in said sample is indicative of the presence of carcinoma in said individual, wherein the absence of TADG-16 in said sample is indicative of the absence of carcinoma in said individual.

The method of claim 38, erein said biological sample is selected from the group consisting of blood, urine, saliva, tears, interstitial

- 40. The method of claim 38, wherein said detection of said TADG-16 is by means selected from the group consisting of Northern blot, Western blot, PCR, dot blot, ELIZA sandwich assay, radioimmunoassay, DNA array chips and flow cytometry of tumor cells, wherein said tumor cells are labeled.
- 41. The method of claim 38, wherein said carcinoma is selected from the group consisting of ovarian, breast, lung, colon, prostate and other in which TADG-16 is overexpressed.
- 42. A method of screening for compounds that inhibit 20 TADG-16, comprising the steps of:
  - (a) contacting a sample with a compound, wherein said sample comprises TADG-16 protein; and

decrease in said TADG-16 protease activity in the presence of said compound relative to TADG-16 protease activity in the absence of said compound is indicative of a compound that inhibits TADG-16.

5

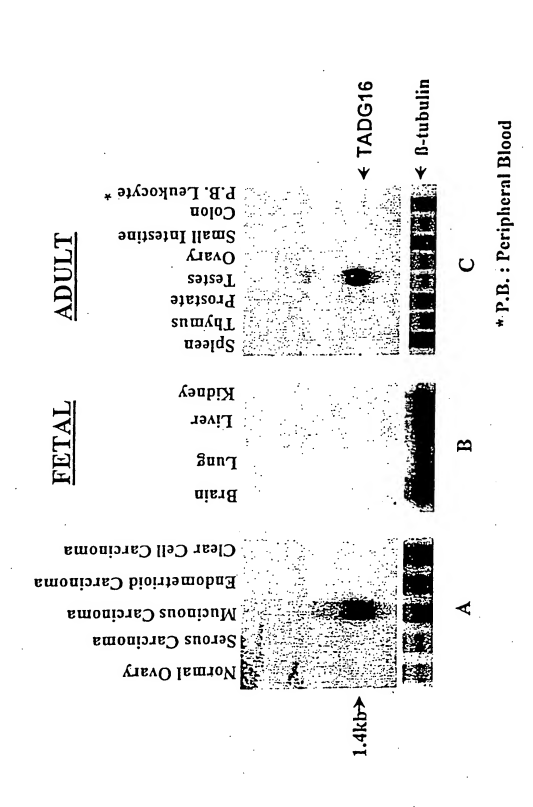
43. A method of targeted therapy to an individual, comprising the step of:

administering a compound to an individual, wherein said 10 compound has a targeting moiety and a therapeutic moiety, wherein said targeting moiety is specific for TADG-16.

- 44. The method of claim 43, wherein said targeting moiety is selected from the group consisting of an antibody specific for TADG-16 and a ligand that binds TADG-16 or a ligand binding domain thereof.
- 45. The method of claim 43, wherein said therapeutic moiety is selected from the group consisting of a radioisotope, a toxin, a chemotherapeutic agent, an immune stimulant and a cytotoxic agent.

The method of claim 43, erein said individual suffers from a cancer selected from the group consisting of ovarian, lung, prostate, colon and others in which TADG-16 is overexpressed.

	1				50	
Prom						
Tryl						
Scce	YYYYMAOKEO	GRTVPCCSRI	P KVAALTAGTI	LLLTAIGAAS	WAIVAVLLRS	
Heps Tadg16	AAAAMAQREG	. ~~~~~~				
raugio				•		
	51				100	
Prom						
Tryl					~~~~~~	
Scce Heps	DOEPT.YPVOV	SSADARLMVE	DKTEGTWRLI	. CSSRSNARVA	GLSCEEMGFL	
Tadg16						
-		•		•	150	
_	101				~~MKKLMVVL	
Prom Tryl				~~~~~~~~	MNPLL.IL	
Scce				~~~~~~MAR	SLLLPLQILL	
Heps	PATTHSELDV	RTAGANGTSG	FFCVDEGRLP	HTQRLLEVIS	VCDCPRGRFL	
Tadg16	~~~~~	~~~~~~		~MGARGALLL	ALLLARAGLR	
	153				200	
Prom	151 SITAAAWA	EEONKLVHG	GPCDKTSHPY	QAALYTSGHL	LCGGVLIHPL	
Tryl	סממזמא מאר מאם	FDDDDDKTVGG	YNCEENSVPY	QVSL.NSGYH	FCGGSLINEQ	
Scce	TOTATEMACE	FACCORTIDG	APCARGSHPW	OVALLSGNOL	HCGGVLVNER	
Heps	AAICQDCGRR	KLPVDRIVGG	RDISLGRWPW	QVSLRYDGAH QGSLRLWDSH	VCGVSLLSHR	
Tadg16	KPTIRGPCGR	KATIZKIAGG	EDALLORMEN	<u> </u>		•
	201	•			250	•
Prom	CHILL WARRICKE	PNLQV	.FLGKHNLRQ	RESS.QEQSS	VVRAVIHPDY	
Tryl	INMICACHOVE	CDIOU	RIGEHNIEV	LEGN. EUFIN	WWITTUIL & 1	
Scce	WVLTAAHCKM	NEYTV	. HLGSDILGD	RRAQRIK ASPH.GLQLG	VOAVVYHGGY	
Heps Tadgl <u>6</u>	WVLTAAHCEP	TYSDLSDPSG	WYVQFGQLTS	MPSFWSLQAY	YTRYFVSNIY	
raddið	JAM TANHOLD					
	251			ODI DI EDDOS	300 A NTTSCHT	
Prom	DAAS	HDODIMLLEL	ARPAKLSELI	STISLPTAPP	ATGTKCLI	
_		CONTINUE OF THE	ハトロルストランドマ	VYAVDEDVCO		
Scce Heps	* DEDDDMCCC	MCMOTAT.VHI.	SSPLPLTEXI	OPVCLPAAGQ	MINDGVICIA	
Tadg16	LSPRYLGNSP	YDIALVKL	SAPVTYTKHI	QPICLQASTF	EFENRTDCWV	
-			•		350	
	301	C DEPOTIC	AYIHLVSREE	CEHA. YPGQ	ITQNMLCAGD	•
Prom Tryl		* PADULIUC	TIDAPVI.SUAK	CEAS. IFGA	TISHITOVOL	
Scce		TO TEDEDIMO	VDVXLISPOD	LIKV. INDL	PENSUIPCUGT	
	TGWGYIKEDE	$\sim \sim \times cut \cap \Gamma$	ADVDIISNDV	CNGADE IGNO	TUE WILL CUG 1	
Tadg16	TGWGYIKEDE	ALPSPHILQE	AGANTIMON	CIMETIZATIO		
	351				400	
Prom		DSGGPLVCGD	HLR	.GLVSWGNIP	CGSKEKPGVY	
Tryl		DECEDITIONS	$\Omega L \Omega$	.GVVSWGD.G	CWOUNTERS	
Scce	PDSKKNACNG PEGGIDACQG	DCCCDEVCED.	SISTIPRWEL	CGIVSWGI.G	CHIMONEGAT	
	PEGGIDACQG	DSGGPLACN.	KNGLWYQ	IGVVSWG.VG	CGRPNRPGVY	
Tadg16						
	401				443	NO 21
Prom	TNVCRYTNWI	QKTIQAK~~~			(SEQ ID	MO: 3)
Tryl	TKVYNYVKWI	KNTIAANS~~			(SEQ ID (SEQ ID	NO: 5)
**	mearcococut	<b>EUVIKARCEV</b>	SCMVTOL~~~	~~~~~~~~	~~~ (SEC ID	NO: 0)
neps Tadg16	TNISHHFEWI	QKLMAQSGMS	QPDPSWPLLF	FPLLWALPLL	GPV (SEQ ID	NO: (7)
910			Fig. 1			



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## Fig. 3A

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301 GACTCCAATC TGATACCACA GTCCATTCTT GTTACAGGCC AAGGGTCCAC

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(SEQ ID NO: 179)

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12	21				AGI S																					acg r
18					rcg . V																					
24					GGG D																	-				
30 ±	1	GC(	GGC A	GCZ (H)	CT	GCT F	TT	GAA E	AC( T	STA Y	TA	GTC	SAC	CT L	TA S	GTG D	ATO	CCC	TC S	CG( G	GGT W	GG:	AT( M	GG1 V	CC C	AG
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1021 1081																										

Fig. 5

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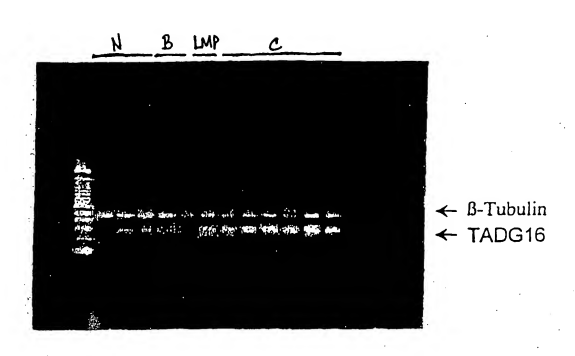


Fig. 6A

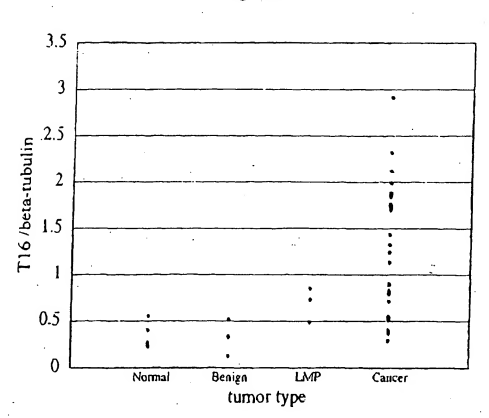


Fig. 6B



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Gly Gly Glu Asp Ala Glu Leu Gly Arg
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Leu Ser Asp Pro Ser Gly Trp Met Val
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Tyr Ser Asp Leu Ser Asp Pro Ser Gly
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Asp Ala Glu Leu Gly Arg Trp Pro Trp
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Leu Leu Leu Ala Leu Leu Ala Arg
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Cys Leu Gln Ala Ser Thr Phe Glu Phe
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His Ile Gln Pro Ile Cys Leu Gln Ala
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Thr Ala Ala His Cys Phe Glu Thr Tyr
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Arg Tyr Phe Val Ser Asn Ile Tyr Leu
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Gly Tyr Ile Lys Glu Asp Glu Ala Leu
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       Residues 201-209 of the TADG-16 protein
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Asn Ser Met Cys Asn His Leu Phe Leu
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		PCT/US00/28558		
A. CLASSIFICATION OF SUBJECT MATTER  1PC(7) : C12N 15/00, 5/00; C12P 21/06; C07H 21/02  US CL : 435/ 320.1, 325, 69.1; 536/ 23.1				
	o International Patent Classification (IPC) or to both DS SEARCHED	national classification and IPC		
Minimum documentation searched (classification system followed by classification symbols) U.S.: 435/320.1, 325, 69.1; 536/23.1				
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) MEDLINE				
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where a		Relevant to claim No.	
X  Y	HOOPER et al., GENBANK (Accession No. AF058300), National Library of Medicine, Bethesda MD., July 1, 1999 (01.04.1999)		1-4, 9-10  6-7,11	
_				
X ·	INOUE, M. et al., Cloning and tissue distribution human eosinophils. Biochem. Biophys. Res. Comm	1-4, 6-10		
X	pages 307-312.  INOUE, M. et al., GENBANK (Accession No. AB031329), National Library of		1-4, 6-10	
 Y	Medicine, Bethesda MD., November 1998, (DNA		11	
x	ONO PHARM CO LTD., GENBANK (Accession No. X15336), National Library of		1-3	
Y	Medicine, Bethesda MD., May 4, 1999 (04.05.199	4-7, 9-11		
x	WO 98/36054 a1 (AMRAD OPERATIONS PTY. see Fig.6, see also pages 14, 33, 38, and 40)	1, 3-7, 9-11		
		·		
Further documents are listed in the continuation of Box C. See patent family annex.				
• S	pecial categories of cited documents:		cument published after the international filing date or priority i not in conflict with the application but cited to understand the	
"A" document defining the general state of the art which is not considered to be of particular relevance		principle or theory underlying the inve	ention	
"E" earlier application or patent published on or after the international filing date "X" document of particular relevance; the claimed invention cannot considered novel or cannot be considered to involve an inventive when the document is taken alone				
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)		"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is		
"O" document	O" document referring to an oral disclosure, use, exhibition or other means  combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents, such combined with one or more other such documents.		•	
"P" document published prior to the international filing date but later than the "&" document member of the same patent family priority date claimed			family .	
Date of the actual completion of the international search  Date of the actual completion of the international search  Date of the actual completion of the international search  Date of the actual completion of the international search				
08 January 2001 (08.01.2001)				
Commissioner of Palents and Trademarks				
Box PCT Washington, D.C. 20231				
	o. (703)305-3230	Telephone No. 703-308-0196	•	

#### INTERNATIONAL SEARCH REPORT

International application No.

PCT/US00/28558

## Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet) This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons: Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely: Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet) This International Searching Authority found multiple inventions in this international application, as follows: Please See Continuation Sheet As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. As only some of the required additional search fees were timely paid by the applicant, this international search 3. report covers only those claims for which fees were paid, specifically claims Nos.: No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-11 Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group 1, claim(s) 1-11, drawn to DNA encoding a TADG-16 protein.

Group 2, claim(s) 12-13, drawn to isolated and purified TADG-16 protein.

Group 3, claim(s) 14, drawn to an antibody.

Group 4, claim(s) 15-20, drawn to a method for detecting TADG-16 mRNA.

Group 5, claim(s) 21-26, drawn to a method for detecting TADG-16 protein.

Group 6, claim(s) 27-28, drawn to a method of inhibiting endogenous expression of TADG-16 in a cell.

Group 7, claim(s) 29-30, drawn to a method of treating a neoplastic state in an individual.

Group 8, claim(s) 31-37, drawn to a method of vaccinating an individual with a TADG-16 fragment or an immunogenic composition consisting of SEQ ID NOs: 17-19, 77-80, 97-99.

Group 9, claim(s) 31-37, drawn to a method of vaccinating an individual with a TADG-16 fragment or an immunogenic composition consisting of SEQ ID NOs: 137-140.

Group 10, claim(s) 31-37, drawn to a method of vaccinating an individual with a TADG-16 fragment or an immunogenic composition consisting of SEQ ID NO: 141.

Group 11, claim(s) 38-41, drawn to a method of diagnosing cancer in an individual.

Group 12, claim(s) 42, drawn to a method of screening for compounds that inhibit TADG-16.

Group 13, claim(s) 43-46, drawn to a method of targeted therapy to an individual.

The inventions listed as Groups 1-11 do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Antalis et al. (WO/9836054, August 1998) teach an isolated DNA encoding a tumor antigen which 100% sequence similarity to SEQ ID NO:2 and or a TADG -16 protein (see Fig 6 and attached sequence comparison).

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